Engineering interfacial structure in “Giant” PbS/CdS quantum dots for photoelectrochemical solar energy conversion

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ARTICLE INFO

Keywords:
Photoelectrochemical
Giant quantum dot
PbS
CdS
Core/shell

ABSTRACT

The interfacial structure in “giant” PbS/CdS quantum dots (QDs) was engineered by modulating the Cd:S molar ratio during in situ growth. The control of the gradient interfacial layer could facilitate hole transfer, regulate the transition from double- to single-color emission, as a consequence. These QDs are optically active close-to-the near-infrared (NIR) spectral region and are candidates as absorber materials in solar energy conversion. Photoinduced charge transfer from “giant” QDs to electron scavenger can still take place despite the ultra-thick (~5 nm) shell. The hybrid architecture based on a TiO2 mesoporous framework sensitized by the “giant” QDs with alloyed interface can produce a saturated photocurrent density as high as ~5.3 mA/cm2 in a photoelectrochemical (PEC) cell under 1 Sun illumination, which is around 2 times higher than that of bare PbS and core/thin-shell PbS/CdS QDs sensitizer. The as-prepared PEC device presented very good stability thanks to the “giant” core/shell QDs architecture with tailored interfacial layer and a further coating of the ZnS shell. 78% of the initial current density is kept after 2-h irradiation at 1 Sun. Engineering of electronic band structure plays a key role in boosting the functional properties of these composite systems, which hold great potential for H2 production in PEC devices.

1. Introduction

Harvesting energy directly from solar radiation provides an attractive approach towards addressing the increasing demand for clean energy, with minimal environmental impacts [1]. Photoelectrochemical (PEC) cells, which directly convert sunlight into electric power or chemical fuels such as hydrogen (H2) are considered a promising route for clean and sustainable energy harvesting [1g,2]. Early studies on photoelectrode materials for PEC cells have been focused on TiO2 and ZnO because of their appropriate energy band position and stability [3]. However, due to their relatively large bandgap of about 3.0–3.2 eV [4], their photoactivation requires ultraviolet (UV)-light, which accounts for only ~5% of the incoming solar energy on the Earth’s surface. To enhance the solar energy conversion, it is necessary to extend the light absorption edge of the photoelectrodes. Semiconductor quantum dot (QD) sensitization has recently been studied intensively thanks to their unique size- and shape-tunable optical properties, including efficient broadband absorption [5,6]. Among various kinds of QDs, lead chalcogenide QDs have large exciton Bohr radius (e.g. 18 nm for PbS) and small bulk energy band gaps (e.g. 0.41 eV for PbS), thus allowing quantum confinement in relatively large sized QDs, together with tunable near-infrared (NIR) absorption and emission [7,8,9], providing potential applications in solar cells [9,10,11] and PEC H2 generation [12,13,14].

Unfortunately, due to their large surface-to-volume ratio, the optical properties of colloidal PbS QDs are very sensitive to their surface chemical conditions (surface ligands, surface oxidation, surface etching, etc.), moisture, oxygen, temperature and/or light [15]. Such
sensitivity contributes to the presence of surface traps/defects, leading to a high density of recombination centers, and hence a considerable decrease of chemical/photo-stability [14,16]. Recent studies revealed that the chemical/photo-stability of conventional colloidal QDs can be improved by post-growth of a robust thick CdS inorganic shell via successive ionic layer adsorption and reaction (SILAR) [12,13]. This thick CdS shell inhibits surface oxidation and the formation of surface traps of PbS core due to better surface passivation and isolation from their environment [17,18]. However, in situ synthesized QDs with thick CdS passivation layer grown via solid-state SILAR suffer from precise control over QD coverage, high charge recombination due to the possible presence of interfacial traps, and is highly time-consuming [12,19,20]. These issues can be addressed by using in-solution pre-synthesized QDs to sensitize the mesoporous films in PEC systems [21]. Size-tunable thick-shell QDs with high quantum yield (QY) and narrow size distribution can be synthesized via an organometallic approach in an organic solvent, introducing surface ligands to passivate the QD surface [22]. A relatively thick shell (in general 3–5 nm thick) [23,24] can efficiently isolate the core material from the QD's surface chemistry and the surrounding chemical environment [23,25,26,27]. However, the sharp interface in type I PbS/CdS core/shell structures may contribute to unwanted confinement of charge carriers and the interfacial defects caused by the 2% lattice mismatch between the rock salt (RS) crystal structure of the core and zinc blende (ZB) structure of the shell [28,29], hindering charge dissociation and transport for solar energy conversion [30,31]. Engineering the electronic band alignment and interfacial structure to balance electron transfer and surface passivation is therefore required. We have recently demonstrated a hole blockade interfacial layer between the core and shell, leading to double color emission in “giant” QDs [22]. This opens the possibility to optimize QD structure to simultaneously maximize carrier transport and surface passivation. By creating a ternary concentration-gradient interfacial layer between the PbS core and the CdS shell, the charge transfer could be facilitated [23,25] due to the partial leakage of the exciton into the shell, as seen for CdS/Te(1−x) [32], CdZn1−xS [33], PbSSe1−x [34], and CdSe/CdZn1−xS [35].

In the present investigation, we report new insight into the synthesis of “giant” PbS/CdS QDs with engineered interfacial layer, demonstrating the possibility to fine tune the hole transfer dynamics by controlling the core/shell interface. Interface modulation leads to a transition from double- to single-color emission, as a consequence of the different charge dynamics between a graded and an abrupt interface. By using high-resolution scanning transmission electron microscopy (HR-STEM), we provide evidence of the presence of Pb in single-emitting QDs. In addition, the photoinduced charge transfer behavior, which is essential for practical exploitation of these systems in PEC devices, was investigated by photoluminescence (PL) and ultrafast pump-probe spectroscopy by monitoring the steady-state PL intensity, PL decay and exciton dynamics in a system composed of “giant” QDs and methyl-viologen chloride (MV2+), an electron scavenger. We performed energy dispersive X-ray spectrometry (EDS) investigations on a Cs-probe-corrected STEM equipped with a high brightness cold field emission gun. In previous work (see Fig. S4 in Ref. [22]), we demonstrated through EDX spectra simulations of a 1 nm PbS core inside a 10 nm CdS QD that only in STEM mode with a 0.1–1 nm sized focused beam, exactly positioned over the particle core, the presence of Pb could be revealed. Experimentally, this beam positioning, as well as the stability of the core and the overall QD under the intense and prolonged electron irradiation, is very critical.

In this work, a particular effort was devoted to finding an experimental demonstration, by STEM techniques, of the presence of a residual PbS core inside the “giant” CdS QDs. To this end, we performed energy dispersive X-ray spectrometry (EDX) investigations using a large area silicon drift detectors (SDD) EDX detector mounted on a Cs-probe-corrected STEM equipped with a high brightness cold field emission gun. In previous work (see Fig. S4 in Ref. [22]), we demonstrated through EDX spectra simulations of a 1 nm PbS core inside a 10 nm CdS QD that only in STEM mode with a 0.1–1 nm sized focused beam, exactly positioned over the particle core, the presence of Pb could be revealed. Experimentally, this beam positioning, as well as the stability of the core and the overall QD under the intense and prolonged electron irradiation, is very critical.

The results of these STEM and EDX combined investigations are summarized in Fig. 1c and d. A high-resolution high angle annular dark field (HAADF) STEM image of a QD (Fig. 1c) clearly shows the crystalline lattice of CdS and a bright triangular central region. As the image contrast in HAADF-STEM also depends on the atomic number (Z-contrast) [42], we used a very high collection inner semi-angle of 130 mrad (β) to enhance the compositional contrast [43] and avoid the coherent effects of elastic scattering almost entirely thermally diffused [44]. This solution, and the possibility to use a high illumination angle of 33 mrad, guaranteed by the presence of the Cs corrector, allowed us to obtain an image with high signal to noise ratio (SNR), where regions of higher intensity may indicate the presence of a heavier QD core, as shown in the bright triangular central region in Fig. 1c. To determine the core composition, an EDX spectrum acquired in a small
square area around the bright triangular central region was recorded (Fig. 1d). In this spectrum, the C, S, Cd and Cu (from the supporting grid) are the main X-ray peaks, however, weak but clear Pb L peaks are detected in this region (inset of Fig. 1d), indicating the presence of Pb in the QD core. It is worth noting, that owing to the critical points previously described, it is not always possible to detect the presence of the Pb L peaks during these experiments. Finally, if the bright central areas visible in QDs HAADF-STEM images like that in Fig. 1c are identified with the PbS core projections, the resulting dimensions of the PbS cores are consistent with the above-reported estimation.

The crystal structure of the QDs was further characterized by X-ray diffraction (XRD, Fig. 1e). The core/shell nanocrystals after cation exchange shift from an RS to a ZB CdS-like pattern [17,22]. After 8 cycles of CdS coating, the overall diffraction pattern exhibits reflections...
from both CdS hexagonal WZ and CdS ZB structure [45,46], irrespective of the Cd:S molar ratio used during the SILAR reaction (Fig. 1e), consistent with HAADF high-resolution STEM investigations (Fig. S4). However, based on the further quantitative analysis (Table S2 and Fig. S5, SI), the relative intensity of peaks related to ZB in double-emitting QDs is almost double compared to that in the single-emitting QDs. This observation is in perfect agreement with the explanation of the double emission as a consequence of the presence of a thick ZB shell acting as a hole-blocking layer [22,45]. No clear reflection from PbS was visible in the XRD pattern of PbS/CdS QDs, even after the growth of a thin CdS shell, due to the very small PbS content (0.1% in vol. in the “giant” system).

The absorption and PL spectra of the QDs with or without SILAR treatment are reported in Fig. 2. The starting bare PbS QDs show a clear and narrow first-exciton peak at 1310 nm, indicating a narrow size distribution, consistent with the TEM observation (Fig. S1d and e). The absorption spectrum of core/thin-shell QDs after cation exchange shifts to lower wavelengths due to the shrinking of the PbS core (Fig. 2a). A CdS molar ratio equal to 1:1 during SILAR results in a double-color emission, with two peaks at 630 nm and 480 nm, respectively (Fig. 2b, blue PL signal). When the Cd:S molar ratio is set to 1:0.8, the “giant” QDs show a typical single emission at 670 nm (Fig. 2b, pink PL signal). As previously reported [17,39], the peak at 480 nm originates from the CdS shell emission, and the one at 630 nm comes from the PbS core. PbS emission has a lifetime in the μs scale, which is typical for PbS QDs [36]. In single-emitting QDs, the band at 670 nm has a lifetime in the μs scale, suggesting the same origin as the double-emitting QDs.

We previously elucidated the mechanism of double-color emitting QDs via pump-probe spectroscopy, which is due to the presence of a core/ZB shell/WZ shell structure (Fig. 1f) [22], in which the ZB shell acts as “hole-blocker” due to a difference in band offsets between ZB and WZ CdS [47]. Now, we can control the interfacial potential barrier between the core and the external shell by modulating the Cd:S molar ratio during SILAR, allowing us to tailor the hole transfer and the double-to-single-color transition in “giant” QDs. When the Cd:S molar ratio is 1:1, at the high reaction temperature, the Cd-oleate reacts with S and does not replace the Pb in the PbS core (cation exchange), forming an abrupt core/shell interface consisting of a PbS core and a CdS ZB shell. At Cd:S molar ratio higher than 1:0.8, the excess Cd still undergoes the cation exchange reaction, which releases free Pb cations. These Pb cations can further react with S to form a Pb$_2$Cd$_{1−x}$S$_x$ alloy, leading to a graded interface (see the scheme in Fig. 1f, right). As shown in Fig. S3, annealing at high temperature (240 °C) in a Cd-oleate-free environment does not induce significant changes in the PL of QDs after cation exchange, suggesting that the PbS core keeps its original size and composition. A clear blue shift is recorded, instead, by dispersing the QDs in Cd-oleate precursors at 240 °C. This behavior actually confirms that, even with a thicker shell around the PbS, the cation exchange still can undergo if an excess of Cd-oleate is present. The PL peak of PbS core in single-emitting “giant” QD presents a red shift compared to that of double-emitting “giant” QDs, indicating the possible existence of alloyed Pb$_2$Cd$_{1−x}$S$_x$ interfacial layer, leading to exciton leakage [45]. The probable presence of the alloyed interfacial layer provides an energy gradient, endowing the fast release of the photogenerated hole from the CdS WZ shell to the PbS core, inhibiting direct exciton radiative recombination within the CdS shell and leading to the single emission of “giant” QDs, while the abrupt core/shell interface leads to double emission [45].

2.2. Charge dynamics of QDs coupled with electron scavenger

To further understand the optical properties of “giant” QDs, we examined their charge transfer behavior by using an electron scavenger, namely MV$_2^{2+}$. The redox energy level of MV$_2^{2+}$ is around 4.06 eV versus the vacuum [31], which is favorable for electron injection from both the PbS and CdS investigated here. The absorption of MV$_2^{2+}$ below 350 nm does not overlap with any PL band of the QDs, excluding any possible energy transfer from the QDs to MV$_2^{2+}$ [31]. The QD sample at the same concentration (details for identifying the concentration are reported in SI and Fig. S2) were mixed with MV$_2^{2+}$, then the charge transfer behavior was investigated by static and transient PL spectra. Taking into account the experimental error, the concentration ratios of MV$_2^{2+}$ and QDs for single- and double-emitting QDs can be regarded as similar. The PL intensity is observed to decrease (Fig. 3a and b) with the increasing concentration of the electron scavengers. Such emission was quenched after the introduction of MV$_2^{2+}$, due to photo-excited charge transfer processes, and the higher quenching rate suggests more efficient electron transfer [31]. To quantitatively study the PL quenching behavior, we considered the PL intensity ratio $q$: 

$$q = 1/I_0$$

where $I_0$ and $I$ are the integrated PL intensities observed without and with quenchers (MV$_2^{2+}$) in solution, respectively. As shown in Fig. 3c, in double-emitting QDs, the PL quenching of the peak attributed to the PbS core is much more intense than the peak attributed to the CdS shell. This is possible due to the rapid radiative recombination (several ns), which occurs in the CdS shell, because charge transfer from the shell cannot effectively compete with the emission process in terms of time scale [17]. On the other hand, the PbS in double-emitting QDs presents more efficient quenching with respect to PbS core in single-emitting QDs (Fig. 3c). This can be explained by examining the structural features of QDs. In case of single-emitting QDs, an alloyed interfacial layer could provide an energy gradient. Once the photo-generated electron of CdS is captured by MV$_2^{2+}$, the photo-generated hole of CdS could quickly migrate to the PbS core to recombine with the electron of PbS, contributing to the core emission. In double-emitting QDs, instead, the ZB shell acting as “hole-blocker” confines the hole in the CdS shell, which can no longer contribute to PbS emission. Moreover, electrons from PbS can tunnel through the CdS barrier, being captured by MV$_2^{2+}$ at the surface of the QDs. To understand the phenomenon, we investigated the fluorescence decay of the “giant” QDs after the introduction of MV$_2^{2+}$, by analyzing the charge transfer rate constants ($k_c$) derived from the fluorescence lifetime. The decay curves of the PL peak corresponding to PbS core and CdS shell, which varied with the concentration of MV$_2^{2+}$, were well fitted with a three-component (PbS emission) and two-component (CdS emission) exponential decay, respectively, shown on a semi-logarithmic plot (Fig. 4a, b and c). To estimate the charge transfer rate constant, the average PL lifetime was calculated from $\tau$ (and $\tau_2$), the characteristic decay times of the two (or three)-component decay. The intensity-weighted average lifetime $<\tau>$ is estimated as follows:
Where $a_i$ are the coefficients of the fitting of PL decay ($i=1, 2$ and $i=1, 2, 3$ for double/tri-exponential decay, respectively). The average PL lifetime of the core largely decreased from around 1330 ns (MV²⁺: QD = 0: 1) to 1040 ns for double-emitting QDs with the addition of MV²⁺ (MV²⁺: QD = 18.9: 1); a similar trend was also found in the single-emitting QDs (Fig. 4a, b). The average PL lifetime of CdS was ~13 ns without the addition of MV²⁺ (Fig. 4c), which is within the range reported for CdS QDs [48]. The charge-transfer rate constant ($k_{et}$) was then estimated using the following equation [49]:

$$k_{et} = \frac{1}{\tau_{QD/MV^{2+}}} - \frac{1}{\tau_{QD}}$$

Where $<\tau_{QD/MV^{2+}}>$ and $<\tau_{QD}>$ are the average PL lifetimes of the QD/MV²⁺ and QDs systems, respectively. Fig. 4d shows the variation of $k_{et}$ with MV²⁺ concentration. The charge-transfer rate of the PbS core significantly increases with the increase of MV²⁺ concentration up to reach a value of $2 \times 10^7$ s⁻¹ (Fig. 4d), which is similar to the $k_{et}$ of single-emitting QDs. On the other hand, as shown in Fig. 4d, the charge transfer rate of CdS emission ($2 \times 10^7$ s⁻¹) is 100 times faster than that of PbS.

In general, photoexcited charge transfer from QDs to charge scavengers relies on geometric factor, time scale and band alignment [17,36,50]. In the "giant" QDs, the PbS QDs are surrounded by the thick CdS shell, which largely slows down the charge transfer from the core. To better understand the role of the shell, we performed pump-probe measurements with a time resolution of about 100 fs collecting the differential transmission $\Delta T/T$ spectra for different pump-probe delays. We adjusted the pump photon energy at the bandgap of CdS so as to disregard exciton thermalization toward the band-edge and focus on exciton recombinaction and charge transfer. We used low pump fluences so as to have an average number of exciton per dot $<N>$ lower than one and avoid multie exciton processes. $\Delta T/T$ spectra (Fig. 5a and b in SI) show a positive signal assigned to bleaching of CdS states, by comparison with the absorption spectrum of the samples (Fig. S6 in SI and Fig. 2b). In fact, the absorption spectra are dominated by the CdS, as PbS contribution cannot be detected due to the high ratio between the volume of the shell and the total volume of "giant" QDs [51], and MV²⁺ shows no absorption in the 300–700 nm range [52]. $\Delta T/T$ signal from CdS, however, reflects the behavior of all excited electrons that are delocalized along the core and the shell, due to a quasi-type-II level alignment of the system [22]. The effect of electron scavenger can be noted in $\Delta T/T$ spectra, which show a reduction of the bleaching signal when MV²⁺ is mixed to QDs (more evident for single-emitting QDs, but also present in "giant" QDs, Fig. S6a and b), while the photoinduced absorption could not be detected due to formation of MV²⁺ radical [31], because it overlaps with photoinduced absorption from defects states. More evidence of the role of the charge scavenger can be observed in $\Delta T/T$ bleaching dynamics (Fig. 5): the presence of the MV²⁺ causes a faster decay of the signal both in the single and double emitting QDs, which is a further unambiguous signature of charge transfer [31].

$\Delta T/T$ bleaching kinetics allows also for an estimation of the charge transfer rate. $\Delta T/T$ in our system reflects the population of photoexcited electrons, whose decay is a consequence of exciton recombinaction in bare QDs samples and of exciton recombinaction and charge transfer in QDs mixed with MV²⁺. We fitted the kinetics and best results were obtained using a bi-exponential function. Using the intensity weighted average lifetime defined above [31,36,53], we obtain a charge transfer rate of about $2 \times 10^7$ s⁻¹ for the single-emitting "giant"
QD and about $3 \times 10^7$ s$^{-1}$ for the double-emitting “giant” QD. These results are consistent with the method proposed by Senty et al. in Ref. [54] for the evaluation of the electron transfer rate behavior as a function of the carrier concentration. We found the same values for time delays longer than few hundreds of picoseconds (while they are one order of magnitude higher for shorter time delays, corresponding to higher carrier concentrations). These results are in agreement with data obtained from PL analyses and show that on the short ns time delays.

![Figure 4](image1.png)

**Fig. 4.** Typical PL decay curves vary without and with the introduction of MV$^{2+}$ for (a) PbS core emission (630 nm) in double-emitting “giant” QDs (b) single-emitting “giant” QDs (670 nm), (c) CdS shell emission (480 nm) in double-emitting “giant” QDs. (d) Charge transfer rate constants derived from the fluorescence lifetime, $k_{et}$, varied with MV$^{2+}$ concentration. All QDs were dispersed in toluene and PL measurements were carried out at ambient temperature with a laser of 444 nm.

![Figure 5](image2.png)

**Fig. 5.** $\Delta T/T$ dynamics for $<N> = 0.4$ for “giant” QDs without (red curves) and with MV$^{2+}$ (blue curves) for (a) single-emitting QDs and (b) double-emitting QDs, at CdS band probe energy (500 nm), together with fit (dotted lines).
scale range electron transfer is mediated by the CdS shell. The charge transfer is still effective (Fig. S6c and d in SI) also in the multie exciton regime.

Due to the passivation of CdS thick shell, the electron transfer rate from the core is largely slowed down, typically much slower than in the pure PbS ($k_e$ of $10^{-7}$ s$^{-1}$) or in thin-shell PbS/CdS QDs/MD $k_{eq}$ of $1-8\times10^{4}$ s$^{-1}$) [17]. With respect to PbS, CdS shows much faster transfer rate, due to the shorter distance from the MD $k_{eq}$, the larger driving force, $\Delta G$ (energy difference between the acceptor and donor systems) and the absence of any additional energy barrier [36]. The less efficient quenching of CdS shell with respect to PbS core in double-emitting “giant” QDs, is possibly due to the rapid radiative recombination (several ns) in CdS shell, because charge transfer from the shell cannot effectively compete with the emission process in times of scale. However, the above results indicate that efficient charge transfer is still possible. By controlling the interfacial structure of “giant” QDs, the hole transfer dynamic could be engineered for the potential use of PbS/CdS “giant” QDs in electronic and optoelectronic devices.

2.3. PEC performance

To demonstrate the potential of “giant” QDs for solar energy conversion applications, they were loaded into TiO$_2$ mesoporous film through EPD [37]. Additional ZnS 2-cycle SILAR deposition was added to deposit a protective coating at QD surface. The PEC activity of the TiO$_2$/QDs system toward the H$_2$ evolution reaction (HER) was studied by employing a three-electrode electrochemical cell configuration [13]. A set of linear-sweep voltammograms with respect to the reversible hydrogen electrode (RHE) ($V_{RHE}=V_{Ag/AgCl}+0.197+pH\times(0.059)$) [12,55] was recorded on different photoanodes in the dark and under simulated solar illumination (AM 1.5 G, 100 mW/cm$^2$). The photo-current density in light gradually grows with the increase in voltage, until a saturated current density is obtained (Fig. 6b). Compared to bare single emitting QD-sensitized photoanode (Fig. S7), the photoanode with ZnS capping layer presents a 3-fold enhancement of the saturated current density, as high as 3.8 mA/cm$^2$ (Fig. 6b), which could be ascribed to the significantly reduced charge recombination by the interposition of an intermediate layer as a ZnS coating [56]. For double-emitting QDs, a saturated current density of 3 mA/cm$^2$ is obtained, which is lower than values obtained for single-emitting QDs. This discrepancy may be understood by considering the possible photoinduced charge transfer, good reproducibility, PbS/CdS “giant” QDs hold wide potential applications in electronics and optoelectronics. Inorganic ligand strategies could help to improve the PEC behavior of “giant” QDs. Further synthesis of “giant” PbS/CdS QDs with emission ranging from 900 to 1200 nm are ongoing to develop a class of QDs optically active in the NIR region for solar energy and optoelectronic applications.

4. Experimental section

4.1. A detailed description of the experimental methods is available in SI

4.1.1. QD synthesis

PbS QDs were synthesized by using OLA as ligands and PbS/CdS thin-shell QDs were synthesized via a two-step cation exchange procedure [60]. Deposition of CdS layer on PbS/CdS QDs followed the procedure described in Dennis et al. [61] For the synthesis of dual-color-emitting “giant” QDs, the molar ratio of Cd/S is equal, while for synthesizing single-emitting “giant” QDs, the molar ratio of Cd/S is around 1:0.8. The QDs were washed by ethanol and then re-dispersed in toluene.

4.1.2. TiO$_2$ film preparation

A thin and compact TiO$_2$ layer was spin coated on FTO-coated glass at 2000 rpm for 60 s by using the commercial solution Ti-Nanoxide BL/SC (Solaronix). Then the films were annealed in air at 500 °C for 30 min after drying and cooled down to room temperature. A blend of active anate particles (~20 nm) and larger anate scatter particles (up to 450 nm) paste (18 NR-AO, paste B, from Dyesol) was tape casted, forming a mesoporous film with thickness ~14 µm, as mea-
Fig. 6. (a) Scheme of the PEC device. Right: The schematic band structure, the “dashed stair” is corresponding to the alloyed interfacial layer in single-emitting QDs and the ZB CdS is located between PbS core and WZ CdS in double-emitting QDs. The arrows indicate the electron and hole transfer process. (b) Photocurrent density versus the applied voltage (vs. RHE) for the TiO2/single-emitting QDs/ZnS and TiO2/double-emitting QDs/ZnS in the dark and under AM 1.5 G illumination at 100 mW/cm². (c) Measured current density of sample TiO2/single-emitting QDs/ZnS (red curve) and TiO2/double-emitting QDs/ZnS (blue curve) as a function of time (normalized) at 0.2 V vs. RHE under 100 mW/cm² illumination with AM 1.5 G filter. Inset table: the current density drop for TiO2/single-emitting QDs/ZnS and TiO2/double-emitting QDs/ZnS after 4000 s. Photocurrent density versus (d) the applied voltage (vs. RHE) and (e) time for sample TiO2/single-emitting QDs without and with CTAB treatment, before applying ZnS SILAR treatment.
sured by contact profilometry.

4.1.3. EPD of the “giant” QDs on the TiO2 film

A pair of TiO2/PtO2 slides vertically immersed in the QDs solution and facing each other with a distance of 1 cm. A voltage of 200 V was applied for 120 min [13]. The samples were then rinsed several times with toluene and dried with N2 at room temperature. After that the anode was immersed in CTAB methanol solution for 1 min and then rinsed by methanol. Later on, two monolayers of ZnS were subsequently deposited by SILAR.

4.1.4. Characterization

Morphology and size distribution (Fig. 1a, b and Fig. S1) of PbS/CdS QDs was characterized by a JEOL 2100F TEM. High-resolution TEM and composition of PbS/CdS single-emitting “giant” QDs was characterized by Cs-probe-corrected JEOL ARM200CF TEM/STEM operated at its full energy of 200 keV, equipped with a cold field emission gun and a 100 mm−2 EDX silicon drift detector (SDD) named “Centurio” (Fig. 1c and d). XRD was performed with a Philips X′pert diffractometer using Cu-Kα radiation source (λ=0.15418 nm). XPS was performed in a VG Escalab 220-XL equipped with a hemispherical analyzer for a Twin Anode X-ray Source. ICP-OES was carried out with Perkin Elmer Model Optima 7300 DV. Absorption spectra were acquired with a Cary 5000 UV–Vis–NIR spectrometer (Varian) with a scan speed of 600 nm/min. Static and transient fluorescence spectroscopy were taken with a Fluorolog®-3 system (Horiba Jobin Yvon). Pump-probe measurements were performed using a Ti:Sapphire laser system (Coherent LIBRA-HE). The PEC performance of the photoelectrodes was evaluated in a typical three-electrode configuration (Coherent LIBRA-HE). The PEC performance of the photoelectrodes was evaluated in a typical three-electrode configuration. The PEC performance of the photoelectrodes was evaluated in a typical three-electrode configuration. The PEC performance of the photoelectrodes was evaluated in a typical three-electrode configuration.


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